## **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 98/56845
COSG 18/66, 18/48, 18/76	A1	(43) International Publication Date: 17 December 1998 (17.12.98)
(21) International Application Number: PCT/US  (22) International Filing Date: 3 June 1998 (  (30) Priority Data. (NYTY) 10 June 1997 (10.06.97)  (71) Applicant 110 tyl GOODRICH COMPANY [US/UK New York New Yorkway, Richfield, OH 44286–936  (72) Inventors N120 t A. Rayi, Ram; 6517 Rosebelle Now New York OH 44039 (US). GOSCEWSKI, Rayin vot. NYT Nast With Street, Lorain, OH 4406  (74) Agents: 11 CKER Mary, Ann et al.; The B.F.Goodri pany 19721 Brecksyille Road, Brecksyille, OH 441 (US).	03.06.9  US]; 402  68 (US)  Avenu , Stanle  55 (US)	CA, CN, CZ, EE, GE, HU, ID, IL, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, RO, RU, SD, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).  Published  With international search report.

## (54) Title: THERMOPLASTIC POLYETHER URETHANE

#### (57) Abstract

A thermoplastic polyether polyurethane having good rebound resilience and low hysteresis properties is prepared by the reaction of a disocyanate with a hydroxyl terminated polyether having a weight average molecular weight of at least 1,400 and a glycol chain extender. The polyether contains alkylene oxide repeat units having from 2 to 6 carbon atoms and the chain extender glycol has from 2 to 16 carbon atoms. The amount of the glycol chain extender is from about 0.25 to about 2 moles per mole of the polyether.

# FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL AM AT AU AZ BA BB BE BF BG	Albania Armenia Austria Australia Azerbaijan Bosnia and Herzegovina Barbados Belgium Burkina Faso Bulgaria	ES FI FR GA GB GE GH GN GR	Spain Finland France Gabon United Kingdom Georgia Ghana Guinea Greece	LS LT LU LV MC MD MG MK	Lesotho Lithuania Luxembourg Latvia Monaco Republic of Moldova Madagascar The former Yugoslav Republic of Macedonia	SI SK SN SZ TD TG TJ TM TR	Slovenia Slovakia Senegal Swaziland Chad Togo Tajikistan Turkmenistan Turkey
BA BB BE BF	Bosnia and Herzegovina Barbados Belgium	GB GE GH GN	United Kingdom Georgia Ghana Guinea	MC MD MG	Monaco Republic of Moldova Madagascar The former Yugoslav	TD TG TJ TM	Chad Togo Tajikistan Turkmenistan

WO 98/56845 PCT/US98/11367

#### THERMOPLASTIC POLYETHER URETHANE

#### FIELD OF THE INVENTION

5

10

15

The present invention relates to thermoplastic polyether urethanes which have very high rebound values and low hysteresis values. More particularly, the present invention relates to such urethanes which are derived from high molecular weight hydroxyl terminated polyether intermediates, and preferably aromatic diisocyanates and aromatic glycol chain extenders.

#### **BACKGROUND OF THE INVENTION**

Heretofore, thermoplastic polyurethanes generally had suitable properties with regard to abrasion resistance, and the like. However, such polyurethanes were generally not suitable in applications wherein the following properties are desired: high melting point, low density, low tensile set, low compression set, and especially good rebound and low hysteresis.

20

25

## SUMMARY OF THE INVENTION

The present invention relates to a polyether urethane having good physical properties which is prepared as by the melt polymerization of a hydroxyl terminated polyether intermediate and a chain extender with a diisocyanate. An important aspect of the invention is that the polyether has a weight average molecular weight such as at least 1,400 with higher molecular weights yielding better results. A large number of different diisocyanates can be utilized and the amount or ratio of the moles of diisocyanate to the moles of hydroxylterminated polyether and chain extender is generally from

10

15

20

25

30

about 0.95 to about 1.05. Another important aspect of the present invention is that the chain extender is a substantially non-branched glycol with aromatic glycols being preferred.

# DETAILED DESCRIPTION OF THE INVENTION

The thermoplastic polyurethanes of the present invention desirably have a hydroxyl terminated polyether intermediate containing alkylene oxide repeat groups of from 2 to 6 carbon atoms, desirably from 2 to 4 carbon atoms, e.g., ethylene oxide or propylene oxide, with 4 carbon atoms, i.e., tetramethylene or butylene oxide being preferred, or combinations thereof. The polyether intermediate desirably is a homopolymer and preferably has a high weight average molecular weight, so that good properties are obtained, of at least about 1,400, desirably at least about 2,000, and preferably at least about 2,500 to about 5,000 or 10,000. Blends of such molecular weight polymers can also be utilized while blends containing low molecular weight components are avoided since the urethane polymer thereof yields poor physicla properties such as hysteresis and rebound. The hydroxyl terminated polyethers which are utilized as the intermediate are well know to the art and literature and are commercially available, as for example, Polymeg 2000 from Quaker Oats or Terethane 2900 from DuPont.

The thermoplastic polyether urethanes of the present invention are generally produced by the so-called one-shot process as, for example, where the hydroxyl terminated polyether intermediate, the chain extender and the diisocyanate are added together, mixed and polymerized. Desirably, the polyether and chain extender are added in one stream and the diisocyanate added in another stream.

While aliphatic diisocyanates can be utilized, aromatic diisocyanates are highly preferred. Moreover, the use of multifunctional isocyanate compounds, i.e., triisocyanates, etc. which cause crosslinking, are avoided and thus the amount thereof is generally less

10

15

20

25

30

than 4 mole percent and preferably less than 2 mole percent based upon the total moles of all of the various isocyanates utilized. The diisocyanates can contain from about 4 to about 20 carbon atoms with about 6 or to 16 being preferred. Examples of aliphatic diisocyanates include hexamethylene diisocyanate, isophorone diisocyanate (IPDI), methylene bis(4-cyclohexylisocyanate), 1,4-cyclohexyl diisocyanate (CHDI) and the like. Examples of preferred aromatic diisocyanates include 1,4-diisocyanatobenzene (PPDI), 1,5-naphthalene diisocyanate (NDI), xylene diisocyanate (XDI), isomers of toluene diisocyanate (TDI), with 4,4'methylenebis(phenylisocyanate), isomers thereof, or oligomers thereof, collectively known as MDI, being especially preferred.

The glycol chain extenders utilized in the present invention are preferably straight chain and crystalline. That is, the weight of any branches of all of the chain extenders utilized based upon the total weight of all chain extenders is generally less than 15 percent and preferably less than 10 percent by weight. Crystallinity helps prevent tackiness and is better achieved when the chain extender has an even number of carbon atoms therein. The chain extenders can be aliphatic with aromatic glycol chain extenders being preferred and contain a total number of carbon atoms of from about 2 to about 16, and preferably from about 6 to about 12. Examples of suitable glycol chain extenders include ethylene glycol, propane glycol, butane glycol, pentane glycol, hexane glycol, benzene glycol, and xylenene glycols which is a mixture of 1,4- di(hydroxymethyl)benzene.

Benzene glycol is preferred and specifically includes hydroquinone, i.e., bis(beta-hydroxyethyl)ether also known as 1,4-di(2-hydroethoxy)benzene; resorcinol, i.e., bis(beta-hydroxyethyl)ether also known as 1,3-di(2-hydroxyethyl)benzene; catechol, i.e., bis(beta-hydroxyethyl)ether also known as 1,2-di(2-hydroxyethoxy)benzene; and combinations thereof. The amount of the glycol chain extender is generally from about 0.25 to about 2.0, desirably from about 0.5 to

10

15

20

25

30

about 1.5, and preferably from about 0.70 to about 1.0 mole per mole of the hydroxyl terminated polyether intermediate.

The mole ratio or amount of the total moles of the one or more different diisocyanates is generally from about 0.95 to about 1.05, and preferably from about 0.98 to about 1.03, based upon the total moles of the one or more hydroxyl terminated polyether intermediates and the one or more chain extenders.

The hydroxyl terminated polyether intermediate, the diisocyanate, and the chain extender, as noted above, are generally added together and reacted in accordance with any conventional urethane reaction method. Preferably, the thermoplastic polyether urethane forming components of the present invention are melt polymerized in the presence of a suitable mixer such as a Banbury, or preferably an extruder. Desirably, the diisocyanate is added separate to the extruder and the polyether intermediate is added as a blend with the chain extender. Suitable processing or polymerization starting temperatures of the disocyanate is from about 100°C to about 200°C, and desirably from about 100° to 150°C and for the blend of the chain extender glycol and polyether intermediate from about 100°C to about 220°C and preferably from about 150° to about 200°C. mixing times in order enable the various components to react and form the thermoplastic polyether urethanes of the present invention are generally from about 2 to about 10 and desirably from about 3 to about 5 minutes.

It is often desirable to utilize catalysts such as stannous and other metal carboxylates as well as tertiary amines. Examples of metal carboxylate catalysts include stannous octoate, dibutyltin dilaurate, phenyl mercuric propionate, lead octoate, iron acetylacetonate, magnesium acetylacetonate, and the like, whereas examples of tertiary amine catalysts include triethylene diamine, and the like. The amount of the one or more catalysts is low, generally from

10

15

20

25

30

about 50 to about 100 parts by weight per million parts by weight of the end polymer formed.

The weight average molecular weight of the polymerized thermoplastic polyether urethanes of the present invention generally range from about 100,000 to about 500,000, desirably from about 150,000 to about 400,000, and preferably from about 175,000 to about 300,000.

The thermoplastic polyether urethanes of the present invention can be mixed with various conventional additives or compounding agents such as fillers, thixotropic agents, extenders, pigments, plasticizers, lubricants, UV absorbers, and the like. Fillers include talc, silicates, clays, calcium carbonate, and the like. The above-noted additives and compounding agents can be used in conventional amounts known to the art and literature.

The thermoplastic polyether urethanes of the present invention have been found to have numerous good and/or unexpected properties. One such property is rebound, for example, as measured by the Bay Shore Rebound Test, i.e., ASTM No. D-2632. Rebound values of at least 60 percent, desirably at least 65 percent, and preferably at least 70 and even 75 percent are achieved. Another favorable property is that the thermoplastic composition has low hysteresis. That is, the hysteresis loss for a 10 percent elongation at room temperature (23°C) is 15 percent or less, generally 12 percent or less, and desirably 9 percent or less, and preferably 6 percent or less. The hysteresis loss at 20 percent elongation at room temperature is 30 percent or less, generally 25 percent or less, desirably 20 percent or less, and preferably 15 percent or less. The thermoplastics also have improved low tensile set (ASTM D-412), i.e., generally 10 percent or less, desirably 8 percent or less, and preferably 6 percent or less. Low compression set values according to ASTM D-395 at 23°C for 22 hours of 25 percent or less, 12 percent or less, and preferably 9 percent or less are obtained. Desirably, the compression set at 70°C for 22 hours is generally 40

10

15

20

25

percent or less, desirably 30 percent or less, and preferably 25 percent or less. Unlike prior polyether urethanes, the urethanes of the present invention according to a differential scanning calorimeter have high melting points, for example, from about 170°C to about 230°C and preferably from about 180°C to about 220°C. The polymers of the present invention also have low densities, for example, specific gravities of 1.10 or less, desirably 1.08 or less and even 1.06 or 1.04 and less. Additionally, the thermoplastic polyether urethane polymers have a Shore D hardness of at least 15 or 20, and even at least 25, 30 or 35 to about 50 or 60.

In addition to the above noted properties, the polyether urethanes of the present invention have favorable properties exhibited by conventional thermoplastic urethanes such as good tear resistance, good abrasion resistance, hydrolytic stability, and the like.

The polymers of the present invention can be utilized wherever such favorable properties are desired. For example, the thermoplastic polyether urethanes of the present invention can be utilized as a solid core in golf balls; as shoe soles; they can be melt spun and used as fibers in hosiery, sports ware, active wear, and in elastic trim such as leg bands and diapers; coil cords such as telephone cords; made into pneumatic tubes; used as profile belts with regard to small motors or for other drive or idler belts; used as recreational wheels such as in-line skate wheels (roller blades), roller skates, or skate boards; and the like.

The invention will be better understood by reference to the following examples which serve to illustrate, but not to limit the scope of the present invention.

#### **EXAMPLES**

Polyester polyol synthesis: All the examples involving polyester polyols were condensation products of 1,4-butane diol (BDO) and adipic acid. These adipate esters were prepared by reacting adipic acid and

10

15

BDO with the overall diol/acid mole ratio being greater than 1.0 and varied depending on the desired molecular weight of the polyol. The reactants were charged to a reactor under dry nitrogen blanket, and held at 125°C under atmospheric pressure until the initial waters of condensation ceased to be evolved. Vacuum was then applied with increasing temperature, and 15 ppm stannous chloride catalysis as necessary, and additional waters of condensation removed until the acid number was reduced below 0.8, whereupon the reaction product was cooled and recovered.

Polymers were prepared by the random melt polymerization method. In this method, the polyol and chain extender are blended together at about 60°C. The blend is then heated to 100-200°C as desired. Diphenylmethane diisocyanate (MDI) is heated separately to 100-150°C as desired, then mixed with the blend. The reactants are vigorously mixed for about 3 minutes, during which time the temperature typically increases to 200-250°C. The polymer melt is discharged into a cooled pan, then cured at 70°C for 1 week after which the polymer is tested for its various properties.

20

#### Example 1

186.83 g of poly (tetramethylene adipate) glycol (PTMAG) with a molecular weight (MW) of 870 (end-group analysis) and 13.17 g of 1.4-butane diol (BDO) were blended together at 110°C to yield a blend molecular weight (BMW) of 555. 90.09 g of 4,4'-diphenylmethane diisocyanate (MDI) was heated to 110°C and then reacted with the blend to give the polymer designated as Example 1.

25

#### Example 2

180.24 g of PTMAG with a MW of 2100 (end-group analysis) and 19.76 g of BDO were blended together at 110°C to yield a BMW of 655. 76.33 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 2.

#### Example 3

168.90 g of PTMAG with a MW of 2159 (end-group analysis) and 31.10 g of benzene glycol (BG) were blended together at 110°C to yield a BMW of 850. 59.04 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 3.

15 <u>Example 4</u>

178.67 g of PTMAG with a MW of 2159 (end-group analysis) and 21.33 g of BG were blended together at 110°C to yield a BMW of 1050. 47.74 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 4.

#### Example 5

185.32 g of PTMAG with a MW of 2159 (end-group analysis)

25 and 14.68 g of BG were blended together at 110°C to yield a BMW of
1250. 40.20 g of MDI was heated to 110°C and then reacted with the
blend to give the polymer designated as Example 5.

### Example 6

.20

119.23 g of poly (tetramethylene ether) glycol (PTMEG) with a MW of 670 (end-group analysis) and 30.76 g of BG were blended together at 120°C to yield a BMW of 450. 60.72 g of MDI was heated to 120°C and then reacted with the blend to give the polymer designated as Example 6.

## Example 7

120 g of PTMEG with a MW of 990 (end-group analysis) and 30 g of BG were blended together at 100°C to yield a BMW of 550.
55.37 g of MDI was heated to 100°C and then reacted with the blend to give the polymer designated as Example 7.

### Example 8

15

20

5

182 29 g of PTMEG with a MW of 1000 (end-group analysis) and 17.71 g of BG were blended together at 110°C to yield a blend molecular weight of 736. 67.93 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 8.

#### Example 9

191.29 g of PTMEG with a MW of 1000 (end-group analysis)
and 8.71 g of BG were blended together at 110°C to yield a blend molecular weight of 850. 58.82 g of MDI was heated to 110°C and then
reacted with the blend to give the polymer designated as Example 9.

25

#### Example 10

268.71 g of PTMEG with a MW of 1040 (end-group analysis) and 31.29 g of BG were blended together at 110°C to yield a BMW of 495. 101.27 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 10.

## Example 11

178.22 g of PTMEG with a MW of 1422 (end-group analysis) and 21.77 g of BG were blended together at 110°C to yield a BMW of 850. 58.82 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 11.

#### Example 12

178.23 g of PTMEG with a MW of 1422 (end-group analysis) and 21.77 g of BG were blended together at 130°C to yield a BMW of 850. 58.93 g of MDI was heated to 130°C and then reacted with the blend to give the polymer designated as Example 12.

# 20 Example 13

200 g of PTMEG with a MW of 1422 (end-group analysis) and 13.91 g of BG were blended together at 110°C to yield a BMW of 1014.28. 52.99 g of MDI was heated to 110°C and then reacted with the blend to give the polymer designated as Example 13.

## Example 14

178.32 g of PTMEG with a MW of 1761 (end-group analysis)
30 and 21.64 g of BG were blended together at 130°C to yield a BMW of
950. 52.71 g of MDI was heated to 130°C and then reacted with the

blend. A drop stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 14.

5

#### Example 15

182.83 g of PTMEG with a MW of 1761 (end-group analysis) and 17.17 g of BG were blended together at 150°C to yield a BMW of 1050. 47.64 g of MDI was heated to 150°C and then reacted with the blend. A drop of stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 15.

### Example 16

15

20

10

277.83 g of PTMEG with a MW of 2025 (end-group analysis) and 22.17 g of BG were blended together at 150°C to yield a BMW of 1204. 62.47 g of MDI was heated to 150°C and then reacted with the blend. A drop of stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 16.

#### Example 17

150 g of poly (propylene ether) glycol (PPG) with a MW of 2226 (end-group analysis) and 11.21 g of BG were blended together at 150°C to yield a BMW of 1300. 31.25 g of MDI was heated to 120°C and then reacted with the blend. A drop of stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 17.

10

15

5

#### Example 18

200 g of PTMEG with a MW of 2953 (end-group analysis) and 11.26 g of BG were blended together at 170°C to yield a BMW of 1696. 31.29 g of MDI was heated to 140°C and then reacted with the blend. A drop of stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 18.

## Example 19

20

25

200 g of PTMEG with a MW of 2953 (end-group analysis) and 12.61 g of BG were blended together at 170°C to yield a BMW of 1618. 33.05 g of MDI was heated to 140°C and then reacted with the blend. A drop of stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 19.

### Example 20

200 g of PTMEG with a MW of 2953 (end-group analysis) and 9.92 g of BG were blended together at 170°C to yield a BMW of 1782. 29.64 g of MDI was heated to 140°C and then reacted with the blend. A drop of stannous octoate (catalyst) was added to the reaction after 2 minutes had elapsed since the start of the reaction. The resulting polymer has been designated as Example 20.

10

15

5

Pertinent information with regard to Examples 1 through 20 are set forth in Table I, along with rebound data. Table II sets forth pertinent information with regard to various physical properties of selected examples. Table III relates to hysteresis data concerning selected examples.

TABLE

ĩ	_		$\top$	T	$\top$	T	7	Ť		Ŧ -			<del></del>	<del></del>			-,		
	Snore		20	35	33	2 6	67	3 3	8	50	30	13	33	30	35	19	37	33	28
٠-	(&) punnage		70	27	37	4	50	24		44	57	56	40	59	63	54	62	29	72
	Temp C		011		110	110	110	120	?	100	110	110	110	110	130	110	150	150	150
Blend	Temp. 'C		110	110	110	110	110	120		100	110	110	110	110	130	110	150	150	150
Blend	Molecular	Weight	555	655	850	1050	1250	450		550	736	850	495	850	850	1014	950	1050	1204
Chain	Extender/	Polyether Mole Ratio	0.68	2.56	2.01	1.30	0.86	0.87		1.25	0.49	0.23	1.35	0.87	0.87	0.49	1.08	0.84	0.84
Chain	Extender		800	BDO	BG	BG	BG	BG		BG	BG	BG	BDO	BG	BG	Bg .	BG	BG	BG
Polyol	Molecular	Weight	870	2100	2159	2159	2159	670		066	1000	1000	1040	1422	1422	1422	1761	1761	2025
Polyol Source		×	BFG	BFG	BFG	BFG	BFG	Quaker Oats	(00) POLYMEG	QO POLYMEG	00 POLYMEG	00 POLYMEG	QO POLYMEG	QO POLYMEG	QO POLYMEG	00 POLYMEG	00 POLYMEG	00 POLYMEG	QO POLYMEG
Polyol Type			Ester	Ester	Ester	Ester	Ester	PTMEG		PTMEG									
Ex. #			-	2	3	4	ည	9		7	8	6	10	11	12	13	14	15	16

$\overline{}$		_			
Shore D	•	19	23	21	21
Rebound (%) Shore D	*	89	77	73	73
MDI Temp. °C		120	140	140	140
Blend Temp. °C		150	170	170	170
Blend Molecular	Weight	1300	1696	1618	1782
Chain Extender/	Polyether Mole Ratio	0.84	0.84	0.94	0.74
Chain		96	BG	86	BG
Polyol Molecular	Weight	2226	2953	2953	2953
Ex. # Polyol Type Polyol Source		Arco R2446	DuPont T2900¹	DuPont T2900	DuPont T2900
Polyol Type		PPG	PTMEG	PTMEG	PTMEG
Ex. #		17	18	19	20

<sup>1</sup> Terethane 2900

_	~~	_		_		·				_												
Fxamole 18	23	1 04	<u>.</u>	4266	27	75.2								:		211	-70	2	77			
Example 16	28	1.06		3599		748		5.6		7.8		22.9		389		189	-69	3	72	i	2330	)
Example 8	30	1.10		3400		710		10		18		61		315		170	-45		57		2480	
Example 10	33	1.12		7000		. 570		13		22		70		475		160	-40		40		4530	
Example 2	33	1.20		6500		550		10		23		65		500		151	-32		37		4000	
Test Method	ASTM D-2240	ASTM D-792		ASTM	D-412/D-638	ASTM	D-412/D-638	ASTM	D-412	ASTM D-395 - 22	hrs @ 23°C	ASTM D-395 - 22	hrs @ 70°C	ASTM D	624, DieC	DSC	DSC		ASTM	D-2632	ASTM	D-790
Properties	Hardness Shore D	Specific Gravity	(a/cc)	Tensile Strength	(psi)	Elongation (%)		Tensile set (%)		Compression set	(%)			Tear Strength	(lb/in)	Melting Point (°C)	Glass Transition	Temp. (°C)	Rebound	Resilience (%)	Flex Modulus (psi)	

- 16 -

SUCCIONATION OUESON I

- 17 -

#### **TABLE III**

#### **Hysteresis**

Example No.	% Hysteresis								
	10% Elongation	20% Elongation	200% Elongation						
8	14.2	26.3	51.0						
16	10.6	24.7	49.3						
18	5.6	11.9	39.7						

Test for Hysteresis: The effect of hysteresis is quantitified as a loss of energy during a deformation cycle. It is measured by subtracting the area under the relaxation curve from the area under the extension curve of a stress-strain diagram and expressing this difference as a percentage of the area located under the extension curve. The stress-strain diagram was developed on an Instron 5500R at a crosshead speed of 2 in/min without any time delay at the end of the extension curve. All samples were of a uniform thickness of 0.075 in.

10

15

20

As apparent from Table I, the thermoplastic urethanes made utilizing an ester intermediate, i.e., Examples 1-5, had poor rebound, even utilizing high molecular weight polyester intermediates and benzene glycol as a chain extender. Thermoplastic urethanes utilizing low molecular weight polyether intermediates outside of the present invention, i.e.; less than 1,400, also generally had poor rebound, i.e., Examples 6-10. However, Examples 11-20 which related to the present invention, had very good rebound properties with higher molecular weight values of the polyether intermediate generally having higher rebound values.

Table II shows that improved physical properties were obtained by thermoplastic polyether urethanes of the present invention, i.e., Examples 16 and 18 as compared to Example 2 wherein a polyester intermediate was utilized, or Examples 8 and 10 wherein low molecular weight polyether intermediates were utilized. Yet, Examples 16 and 18 still retained other thermoplastic properties such as good tear strength.

Table III shows good low hysteresis values were obtained utilizing the thermoplastic polyether urethanes of the present invention.

While in accordance with the Patent Statutes, the best mode and preferred embodiment has been set forth, the scope of the invention is not limited thereto, but rather by the scope of the attached claims.

10

15

20

30

#### WHAT IS CLAIMED IS:

1. A thermoplastic polymer, comprising;

a polyether urethane derived from a diisocyanate reacted with a hydroxyl terminated polyether and a glycol chain extender,

said hydroxyl terminated polyether having alkylene oxide repeat units containing from 2 to 6 carbon atoms and having a weight average molecular weight of at least 1,400,

said chain extender being a substantially non-branched glycol having from 2 to 16 carbon atoms, and wherein the amount of said chain extender is from about 0.25 to about 2 moles per mole of said hydroxyl terminated polyether.

- 2. A thermoplastic polymer according to Claim 1, wherein said polyether has alkylene oxide repeat units containing from 2 to 4 carbon atoms.
  - 3. A thermoplastic polymer according to Claim 2, wherein said disocyanate has a total of from 4 to 20 carbon atoms, and wherein the mole ratio of said disocyanate to said hydroxyl terminated polyether and said glycol chain extender is from about 0.95 to about 1.05.
- 4. A thermoplastic polymer according to Claim 3, wherein the weight of any branches in said glycol chain extender is 15 percent or less based upon the total weight of all said chain extenders, wherein said chain extender is an aromatic having a total of from 6 to 12 carbon atoms, and wherein said diisocyanate is an aromatic diisocyanate having a total of from 8 to 16 carbon atoms.

5. A thermoplastic polymer according to Claim 4, wherein said hydroxyl terminated polyether has a weight average molecular

10

weight of at least 2,500, wherein said polyether is polytetramethylene glycol, wherein the mole ratio of said chain extender to said hydroxyl terminated polyether is from about 0.5 to about 1.5, and wherein said thermoplastic polymer has a weight average molecular weight of from about 100,000 to about 500,000.

- 6. A thermoplastic polymer according to Claim 5, wherein said polytetramethylene glycol has a molecular weight of from about 2,500 to about 10,000, wherein the mole ratio of said chain extender to said hydroxyl terminated polyether is from about 0.7 to about 1.0, wherein said chain extender is benzene glycol, and wherein said diisocyanate is MDI.
- 7. A thermoplastic polymer according to Claim 1, wherein 15 said polymer has a rebound of at least 60 percent.
  - 8. A thermoplastic polymer according to Claim 4, wherein said polymer has a rebound of at least 65 percent.
- 9. A thermoplastic polymer according to Claim 6, wherein said polymer has a rebound of at least 70 percent.
- 10. A thermoplastic polymer according to Claim 1, wherein the hysteresis of said polymer at 10 percent elongation is 15 percent or less.
  - 11. A thermoplastic polymer according to Claim 4, wherein the hysteresis of said polymer at 10 percent elongation is 12 percent or less.

- 12. A thermoplastic polymer according to Claim 6, wherein the hysteresis of said polymer at 10 percent elongation is 6 percent or less.
- 13. A thermoplastic polymer according to Claim 1, wherein said polymer has a rebound of at least 60 percent, and wherein the hysteresis of said polymer at 10 percent elongation is 15 percent or less.
- 10 14. A thermoplastic polymer according to Claim 5, wherein said polymer has a rebound of at least 70 percent, and wherein the hysteresis of said polymer at 10 percent elongation is 9 percent or less.
- 15. A golf ball core, comprising; the thermoplastic polymer of Claim 1.
  - 16. A golf ball core, comprising; the thermoplastic polymer of Claim 5.
  - 17. A cord, comprising; the thermoplastic polymer of Claim 1.
- 18. A cord, comprising; the thermoplastic polymer of 25 Claim 6.
  - 19. A recreational wheel, comprising; the thermoplastic polymer of Claim 1.
- 20. A recreational wheel, comprising; the thermoplastic polymer of Claim 6.

- 21. A profile belt, comprising; the thermoplastic polymer of Claim 1.
- 22. A profile belt, comprising; the thermoplastic polymerof Claim 6.
  - 23. An article containing fibers, comprising; the thermoplastic polymer of Claim 1.
- 10 24. An article containing fibers, comprising; the thermoplastic polymer of Claim 6.
  - 25. A process for producing a thermoplastic polyether urethane comprising, the steps of;
- mixing and reacting a hydroxyl terminated polyether, a glycol chain extender, and a diisocyanate, said hydroxyl terminated polyether having alkylene oxide repeat units containing from 2 to 6 carbon atoms and having a weight average molecular weight of at least 1,400.

25

- 26. A process according to Claim 25, wherein said polyether has alkylene oxide repeat units containing from 2 to 4 carbon atoms, wherein said diisocyanate has a total of from 4 to 20 carbon atoms, and wherein the mole ratio of said diisocyanate to said hydroxyl terminated polyether and said glycol chain extender is from about 0.95 to about 1.05.
- 27. A process according to Claim 26, wherein said chain extender is an aromatic glycol having a total of from 6 to 12 carbon atoms, wherein said diisocyanate is an aromatic diisocyanate having a total of from 8 to 16 carbon atoms, and wherein the mole ratio of said chain extender to said polyether is from about 0.5 to about 1.5.

28. A process according to Claim 27, wherein the weight average molecular weight of said hydroxyl terminated polyether is at least 2,500, wherein said polyether is polytetramethylene glycol, wherein said chain extender is benzene glycol, and wherein said diisocyanate is MDI.

## INTERNATIONAL SEARCH REPORT

Inters. Just Application No PCT/US 98/11367

		1017 00 307	11307
IPC 6	IFICATION OF SUBJECT MATTER C08G18/66 C08G18/48 C08G18/	76	
According t	to International Patent Classification(IPC) or to both national classific	ation and IPC	
	SEARCHED		
	ocumentation searched (classification system followed by classificat	ion symbols)	
IPC 6	C08G		•• .
Documenta	ation searched other than minimum documentation to the extent that s	such documents are included in the fields sear	ched
Electronic o	data base consulted during the international search (name of data ba	ase and, where practical, search terms used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category °	Citation of document, with indication, where appropriate, of the rel	evant passages	Relevant to claim No.
X	US 5 545 706 A (N.BARKSBY ET AL. August 1996 see column 11, line 38 - column		1-9
	32; claims 1,11	12, Tine	
Α	US 4 120 850 A (E.PECHHOLD) 17 0	ctober	1-9,21, 22
	see column 2, line 64 - line 68; 1,2; table I	claims	22
Α	EP 0 343 985 A (E.I.DU PONT DE N CO.) 29 November 1989	EMOURS &	1-5, 10-12,
	see page 2, line 16 - line 30		23,24
	·	·	
Furt	her documents are listed in the continuation of box C.	X Patent family members are listed in	annex.
° Special ca	ategories of cited documents:	"T" lotor decompost multiplicate de file al la	
consid	ent defining the general state of the art which is not dered to be of particular relevance	"T" later document published after the internal or priority date and not in conflict with the cited to understand the principle or theolinvention	e application but
· ming c		"X" document of particular relevance; the cla cannot be considered novel or cannot b	Imed invention
citation	ent which may throw doubts on priority claim(s) or is cited to establish the publicationdate of another n or other special reason (as specified)	"Y". document of particular relevance; the cla	ment is taken alone imed invention
otneri	ent referring to an oral disclosure, use, exhibition or means ent published prior to the international filing date but	cannot be considered to involve an inve document is combined with one or more ments, such combination being obvious in the art.	other such docu-
later tr	nan the priority date claimed	"&" document member of the same patent fa	
	actual completion of theinternational search	Date of mailing of the international search	h report
<del></del>	September 1998	18/09/1998	· ·
Name and r	nailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	Authorized officer	
	Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Angiolini, D	

## INTERNATIONAL SEARCH REPORT

information on patent family members

PCT/US 98/11367

Patent document sated in search report	_	Publication date		tent family ember(s)	Publication date
US 5545706	A	13-08-1996	AU AU BR CA CN EP JP	694644 B 5215296 A 9602159 A 2175402 A 1146464 A 0742240 A 8301966 A	23-07-1998 21-11-1996 30-12-1997 10-11-1996 02-04-1997 13-11-1996 19-11-1996
US 4120850	A	17-10-1978		870204 A 1136334 A 2832588 A 2401942 A 2003898 A,B 1347143 C 54050100 A 58007648 B 7809072 A	05-03-1979 23-11-1982 08-03-1979 30-03-1979 21-03-1979 13-11-1986 19-04-1979 10-02-1983 08-03-1979
EP 343985	Α	29-11-1989		5000899 A 1318048 A 1038318 A 68917212 D 68917212 T 2019511 A 9707430 B 171229 B	19-03-1991 18-05-1993 27-12-1989 08-09-1994 16-02-1995 23-01-1990 08-05-1997 12-10-1993